```
? ds
Set
       Items
               Description
        3643
               SARS OR HSARS OR SCOV
S1
      270156
S2
               POLYMERASE
S3
         309
               S1 AND S2
               S3 NOT (PCR OR REVERSE)
S4
         70
S5
        2455 REPLICASE
         50 S1 AND S5
S6
S7
               ORF1 OR ORF1A OR ORF1AB
        1236
S8
           7
               S1 AND S7
       76283
               ANTIBODY AND RESPONSE
S9
S10
         51
               S1 AND S9
S11
        8032
               POL
S12
          11
               S10 AND (S11 OR S2 OR S5 OR S7)
? log hold
      06mar06 12:29:18 User208669 Session D2995.2
           $9.87 2.903 DialUnits File155
              $0.00 148 Type(s) in Format 6
              $2.42 11 Type(s) in Format 7
           $2.42 159 Types
   $12.29 Estimated cost File155
    $3.73 TELNET
   $16.02 Estimated cost this search
   $16.44 Estimated total session cost 3.024 DialUnits
Logoff: level 05.10.03 D 12:29:18
```

Record Date Completed: 20031110

OM nucleic - nucleic search, using sw model

Run on: July 20, 2005, 17:03:25; Search time 78.378 Seconds

(without alignments)

438.412 Million cell updates/sec

Title: US-10-848-737-4

Perfect score: 21

Sequence: 1 gcacuugucuaccuugaugtt 21

Scoring table: IDENTITY NUC

Gapop 10.0 , Gapext 1.0

Searched: 1202784 seqs, 818138359 residues

Total number of hits satisfying chosen parameters: 2405568

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 300 summaries

1: /cgn2\_6/ptodata/1/ina/5A COMB.seq:\*

2: /cgn2\_6/ptodata/1/ina/5B COMB.seq:\*

3: /cgn2\_6/ptodata/1/ina/6A\_COMB.seq:\*

4: /cgn2\_6/ptodata/1/ina/6B\_COMB.seq:\*

5: /cgn2\_6/ptodata/1/ina/PCTUS\_COMB.seq:\*

6: /cgn2\_6/ptodata/1/ina/backfiles1.seq:\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

			ક				
Result			Query				
	No.	Score	Match	Length	DB	ID	Description
С	1	17.4	82.9	393	3	US-09-404-879A-374	Sequence 374, App
С	2	17.4	82.9	393	4	US-09-667-857-374	Sequence 374, App
С	3	17.4	82.9	1463	4	US-09-949-016-2910	Sequence 2910, Ap
C	4	17.4	82.9	1520	4	US-09-919-039-342	Sequence 342, App
C	5	17.4	82.9	1538	4	US-09-976-594-52	Sequence 52, Appl
C	6	17.4	82.9	1542	4	US-09-919-039-341	Sequence 341, App
C	7	17.4	82.9	1990	4	US-09-976-594-53	Sequence 53, Appl
C	8	17	81.0	74730	4	US-09-949-016-15189	Sequence 15189, A
C	9	16.8	80.0	924	4	US-09-248-796A-844	Sequence 844, App
	10	16.8	80.0	1099	4	US-08-956-171E-825	Sequence 825, App
	11	16.8	80.0	1099	4	US-08-781-986A-825	Sequence 825, App

OM nucleic - nucleic search, using sw model

Run on: July 20, 2005, 17:34:20 ; Search time 428.102 Seconds

(without alignments)

311.209 Million cell updates/sec

Title: US-10-848-737-4

Perfect score: 21

Sequence: 1 gcacuugucuaccuugaugtt 21

Scoring table: IDENTITY NUC

Gapop 10.0 , Gapext 1.0

Searched: 7173243 segs, 3172129809 residues

Total number of hits satisfying chosen parameters: 14346486

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 300 summaries

Database : Published Applications NA:\*

/cgn2\_6/ptodata/2/pubpna/US07\_PUBCOMB.seq:\*

2: /cgn2 6/ptodata/2/pubpna/PCT NEW PUB.seq:\*

/cgn2 6/ptodata/2/pubpna/US06 NEW PUB.seq:\* 3:

/cgn2\_6/ptodata/2/pubpna/US06\_PUBCOMB.seq:\*

/cgn2\_6/ptodata/2/pubpna/US07 NEW PUB.seq:\*

6: /cgn2\_6/ptodata/2/pubpna/PCTUS PUBCOMB.seq:\*

7: /cgn2\_6/ptodata/2/pubpna/US08 NEW PUB.seq:\*

/cgn2\_6/ptodata/2/pubpna/US08\_PUBCOMB.seq:\*

/cgn2\_6/ptodata/2/pubpna/US09A\_PUBCOMB.seq:\* 9:

/cgn2\_6/ptodata/2/pubpna/US09B PUBCOMB.seq:\* 10:

11: /cgn2\_6/ptodata/2/pubpna/US09C PUBCOMB.seg:\*

12: /cgn2\_6/ptodata/2/pubpna/US09 NEW PUB.seq:\*

13: /cgn2\_6/ptodata/2/pubpna/US10A PUBCOMB.seq:\*

/cgn2\_6/ptodata/2/pubpna/US10B\_PUBCOMB.seq:\*

15: /cgn2 6/ptodata/2/pubpna/US10C PUBCOMB.seg:\*

/cgn2\_6/ptodata/2/pubpna/US10D\_PUBCOMB.seq:\* 16:

/cgn2 6/ptodata/2/pubpna/US10E\_PUBCOMB.seq:\* 17:

18: /cgn2\_6/ptodata/2/pubpna/US10F PUBCOMB.seq:\*

19: /cgn2\_6/ptodata/2/pubpna/US10G PUBCOMB.seq:\*

20: /cgn2 6/ptodata/2/pubpna/US10H PUBCOMB.seg:\*

/cgn2 6/ptodata/2/pubpna/US10I PUBCOMB.seq:\* 21:

22: /cgn2 6/ptodata/2/pubpna/US10 NEW PUB.seq:\*

23: /cgn2 6/ptodata/2/pubpna/US11A PUBCOMB.seq:\* 24: /cgn2\_6/ptodata/2/pubpna/US11\_NEW\_PUB.seq:\*

25: /cgn2\_6/ptodata/2/pubpna/US60 NEW PUB.seq:\*

26: /cgn2\_6/ptodata/2/pubpna/US60 PUBCOMB.seq:\*

OM nucleic - nucleic search, using sw model

Run on: July 20, 2005, 16:07:29; Search time 271.346 Seconds

(without alignments)

458.140 Million cell updates/sec

Title: US-10-848-737-4

Perfect score: 21

Sequence: 1 gcacuugucuaccuugaugtt 21

Scoring table: IDENTITY NUC

Gapop 10.0 , Gapext 1.0

4390206 seqs, 2959870667 residues Searched:

Total number of hits satisfying chosen parameters: 8780412

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 300 summaries

Database : N Geneseg 16Dec04:\*

1: genesegn1980s:\*

2: geneseqn1990s:\*

3: geneseqn2000s:\*

4: geneseqn2001as:\*

5: geneseqn2001bs:\*

6: geneseqn2002as:\*

7: geneseqn2002bs:\*

8: geneseqn2003as:\*

9: geneseqn2003bs:\*

10: geneseqn2003cs:\*

11: geneseqn2003ds:\*

12: geneseqn2004as:\* 13: geneseqn2004bs:\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

Re	sult No.	Score	Query Match	Length	DB	ID	Description
	1	19	90.5	29751	12	ADJ39000	Adj39000 SARS coro
	2	17.8	84.8	237961	6	ABQ80552	Abq80552 Human Can
C	3	17.4	82.9	355	10	ACD93215	Acd93215 Human col
C	4	17.4	82.9	393	3	AAA70063	Aaa70063 Human ova

OM nucleic - nucleic search, using sw model

July 20, 2005, 16:07:59 ; Search time 496.063 Seconds Run on:

(without alignments)

2051.271 Million cell updates/sec

Title: US-10-848-737-4

Perfect score: 21

Sequence: 1 gcacuugucuaccuugaugtt 21

Scoring table: IDENTITY NUC

Gapop 10.0 , Gapext 1.0

Searched: 4708233 seqs, 24227607955 residues

Total number of hits satisfying chosen parameters: 9416466

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 300 summaries

Database : GenEmbl:\*

1: qb ba:\*

2: gb htg:\*

3: gb in:\*

4: gb om:\*

5: gb\_ov:\*

6: gb\_pat:\*

7: gb ph:\*

8: gb pl:\*

9: gb\_pr:\*

10: gb\_ro:\*

11: gb\_sts:\*

12: gb\_sy:\* 13: gb\_un:\*

14: gb\_vi:\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

Result No.	Score	% Query Match	Length	DB	ID	Description
1	19	90.5	1706	14	AY322205S1	AY322205 SARS coro
2	19	90.5	6200	14	AY534762S1	AY534762 SARS coro
3	19	90.5	14011	14	AY534758S1	AY534758 SARS coro

OM nucleic - nucleic search, using sw model

Run on: June 23, 2005, 23:08:06; Search time 728.716 Seconds

(without alignments)

1396.372 Million cell updates/sec

Title: US-10-848-737-4

Perfect score: 21

Sequence: 1 gcacuugucuaccuugaugtt 21

Scoring table: IDENTITY NUC

Gapop 10.0 , Gapext 1.0

Searched: 4708233 seqs, 24227607955 residues

Total number of hits satisfying chosen parameters: 9416466

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 45 summaries

Database : GenEmbl:\*

1: gb ba:\*

2: gb htg:\*

3: gb\_in:\*

4: gb\_om:\*

5: gb\_ov:\*

6: gb pat:\*

7: gb\_ph:\*

8: gb\_pl:\*

9: gb\_pr:\*

10: gb\_ro:\*

11: gb sts:\*

12: gb\_sy:\*

13: gb un:\*

14: gb\_vi:\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

Result No.	Score	% Query Match	Length	DB	ID	Description	
1	19	90.5	 1706	14	AY322205S1	AY322205 SARS CO	- -
2	19	90.5			AY534762S1	AY534762 SARS CO	
3	19	90.5	14011	14	AY534758S1	AY534758 SARS co	ro

OM nucleic - nucleic search, using sw model

Run on: July 20, 2005, 16:45:00; Search time 2427.07 Seconds

(without alignments)

329.347 Million cell updates/sec

Title: US-10-848-737-4

Perfect score: 21

Sequence: 1 gcacuugucuaccuugaugtt 21

Scoring table: IDENTITY NUC

Gapop 10.0 , Gapext 1.0

Searched: 34239544 seqs, 19032134700 residues

Total number of hits satisfying chosen parameters: 68479088

Minimum DB seq length: 0

Maximum DB seq length: 2000000000

Post-processing: Minimum Match 0%

Maximum Match 100%

Listing first 300 summaries

Database : EST:\*

1: gb\_est1:\*

2: gb est2:\*

3: gb htc:\*

4: gb\_est3:\*

5: gb\_est4:\*

6: gb\_est5:\*

7: gb est6:\*

8: gb\_gss1:\*

9: gb\_gss2:\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

Res	ult		Query					
	No.	Score	Match	Length	DB	ID	Description	
	1	19	90.5	449	2	BE938410	BE938410 RC0-TN007	,
С	2	18.4	87.6	1076	4	BM459897	BM459897 AGENCOURT	•
C	3	18.4	87.6	1128	4	BG110924	BG110924 602284573	į
C	4	18	.85.7	587	5	BP284256	BP284256 BP284256	
C	5	17.8	84.8	342	9	CE508857	CE508857 tigr-gss-	
C	6	.17.8	84.8	600	2	BF356298	BF356298 QV4-HT089	)
	7	17.8	84.8	634	6	CB082521	CB082521 hn52c01.c	ſ
С	8	17.8	84.8	702	5	BW403702	BW403702 BW403702	